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				500,000 in Key STN Databases
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NEWS	12	APR	02	New Thesaurus Added to Derwent Databases for Smooth
				Sailing through U.S. Patent Codes
NEWS	13	APR	02	EMBASE Adds Unique Records from MEDLINE, Expanding
				Coverage back to 1948
NEWS	14	APR	0.7	CA/CAplus CLASS Display Streamlined with Removal of
110110	1.5		0.77	Pre-IPC 8 Data Fields
NEWS	15	APR	U /	50,000 World Traditional Medicine (WTM) Patents Now
NEWS	10	3.00	0.7	Available in CAplus
NEWS	ТР	APR	U /	MEDLINE Coverage Is Extended Back to 1947
MEMC	FVDI	DECC	FFDI	RUARY 15 10 CURRENT WINDOWS VERSION IS V8.4.2,
MEMO	EAL	KEUU		CURRENT DISCOVER FILE IS DATED 15 JANUARY 2010.
				COLLEGE DECOURTE ELECTION TO CHROMIC BOLD.
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\* \*

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=> FILE REGISTRY

COST IN U.S. DOLLARS FULL ESTIMATED COST

SINCE FILE TOTAL SESSION ENTRY 0.22 0.22

FILE 'REGISTRY' ENTERED AT 10:06:35 ON 04 JUN 2010 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS.

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STRUCTURE FILE UPDATES: 2 JUN 2010 HIGHEST RN 1226851-61-1 DICTIONARY FILE UPDATES: 2 JUN 2010 HIGHEST RN 1226851-61-1

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Uploading C:\Program Files\Stnexp\Oueries\10591921z.str

```
17 18 19 20 21 23 24 25 26 27 33 36 ring nodes:
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 22 28 29 30 31 32 chain bonds:
2 -17 5-24 5-25 6-18 7-19 8-20 9-26 13-23 16-21 21-27 21-36 27-28 31-33 ring bonds:
1 -2 1-16 2-3 3-4 4-5 5-6 6-7 7-8 8-9 9-10 10-11 11-12 12-13 13-14 13-22 14-15 14-22 15-16 28-29 28-32 29-30 30-31 31-32 exact home bonds:
2 -17 6-18 8-20 13-23 28-32 31-32 exact bonds:
2 -17 16-18 8-20 13-23 28-32 31-32 exact bonds:
3 -2 14-15 14-22 15-16 28-29 28-32 29-30 30-31 31-32 exact home signature for the first state of the first st
```

G1:H,Ak

Match level :

chain nodes :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:CLASS 18:CLASS 19:CLASS 20:CLASS 21:CLASS 22:Atom 23:CLASS 24:CLASS 25:CLASS 26:CLASS 27:CLASS 28:Atom 29:Atom 30:Atom 31:Atom 33:CLASS 36:CLASS 36:CLASS

### Stereo Bonds:

```
19-7 (Single Hash).
20-8 (Single Wedge).
21-16 (Single Wedge).
26-9 (Single Hash).
```

### Stereo Chiral Centers:

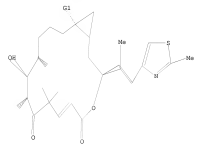
```
7 (Parity=Even)
8 (Parity=Odd)
9 (Parity=Odd)
16 (Parity=Even)
```

### Stereo RSS Sets:

Type=Relative (Default). 4 Nodes= 7 8 9 16

### L1 STRUCTURE UPLOADED

=> d l1 L1 HAS NO ANSWERS L1 STR



G1 H,Ak

Structure attributes must be viewed using STN Express query preparation.

=> s l1

SAMPLE SEARCH INITIATED 10:06:59 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 45 TO ITERATE

100.0% PROCESSED 45 ITERATIONS 0 ANSWERS

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

PROJECTED ITERATIONS:

BATCH \*\*COMPLETE\*\*
498 TO 1302

PROJECTED ITERATIONS: 498 TO 130: PROJECTED ANSWERS: 0 TO

L2 0 SEA SSS SAM L1

=> s 11 sss full FULL SEARCH INITIATED 10:07:07 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 897 TO ITERATE

100.0% PROCESSED 897 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.01

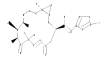
SEARCH TIME: 00.00.01

L3 0 SEA SSS FUL L1

\_

Uploading C:\Program Files\Stnexp\Queries\10591921z1.str





```
ring bonds :
1-2 1-16 2-3 3-4 4-5 5-6 6-7 7-8 8-9 9-10 10-11 11-12 12-13 13-14
13-22 14-15 14-22 15-16 28-29 28-32 29-30 30-31 31-32
exact/norm bonds :
1-2 1-16 2-3 2-17 3-4 4-5 5-6 5-24 5-25 6-7 6-18 7-8 7-19 8-9 8-20
9-10 9-26 10-11 11-12 12-13 13-14 13-22 13-23 14-15 14-22 15-16 16-21
21-27 21-36 27-28 28-29 28-32 29-30 30-31 31-32 31-33
isolated ring systems :
containing 1 : 28 :
G1:H,Ak
G2:C,O
Match level :
```

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:CLASS 18:CLASS 19:CLASS 20:CLASS 21:CLASS 22:Atom 23:CLASS 24:CLASS 25:CLASS 26:CLASS 27:CLASS

28:Atom 29:Atom 30:Atom 31:Atom 32:Atom 33:CLASS 36:CLASS

Stereo Bonds: 19-7 (Single Hash). 20-8 (Single Wedge). 21-16 (Single Wedge).

# 26-9 (Single Hash). Stereo Chiral Centers:

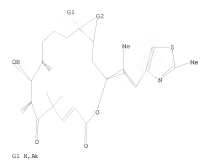
(Parity=Even) 8 (Parity=Odd) 9 (Parity=Odd) 16 (Parity=Even)

### Stereo RSS Sets:

Type=Relative (Default). 4 Nodes= 7 8 9 16

#### L4 STRUCTURE UPLOADED

=> d 14 L4 HAS NO ANSWERS T. 4 STR G2 C, O



Structure attributes must be viewed using STN Express query preparation.

=> s 14 SAMPLE SEARCH INITIATED 10:09:35 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 45 TO ITERATE

100.0% PROCESSED 45 ITERATIONS 0 ANSWERS SEARCH TIME: 00.00.02

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*
BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 498 TO 1302 PROJECTED ANSWERS: 0 TO 0

L5 0 SEA SSS SAM L4

=> s 14 sss full FULL SEARCH INITIATED 10:09:47 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 779 TO ITERATE

100.0% PROCESSED 779 ITERATIONS 6 ANSWERS

SEARCH TIME: 00.00.01

L6 6 SEA SSS FUL L4

=> FIL HCAPLUS COST IN U.S. DOLLARS

 COST IN U.S. DOLLARS
 SINCE FILE
 TOTAL

 FULL ESTIMATED COST
 385.00
 385.00

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FILE COVERS 1907 - 4 Jun 2010 VOL 152 ISS 24 FILE LAST UPDATED: 3 Jun 2010 (20100603/ED) REVISED CLASS FIELDS (/NCL) LAST RELOADED: Apr 2010 USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Apr 2010

HCAplus now includes complete International Patent Classification (IPC) reclassification data for the second quarter of 2010.

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http://www.cas.org/legal/infopolicy.html

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 16 L7 19 L6

=> s 17 and py<=2004 25158292 PY<=2004 17 L7 AND PY<=2004

=> d 17 ibib abs hitstr tot

L7 ANSWER 1 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 2008:1383637 HCAPLUS

DOCUMENT NUMBER: 149:555127

TITLE: Dioxirane epoxidation of alkenes

AUTHOR(S): Adam, Waldemar; Saha-Moeller, Chantu R.; Zhao,

Cong-Gui

CORPORATE SOURCE: Universitaet Wuerzburg, Wuerzburg, Germany

SOURCE: Organic Reactions (Hoboken, NJ, United States) (2002),

61, No pp. given

CODEN: ORHNBA

URL: http://www3.interscience.wilev.com/cgi-

bin/mrwhome/107610747/HOME PUBLISHER: John Wiley & Sons, Inc.

DOCUMENT TYPE: Journal; General Review; (online computer file)

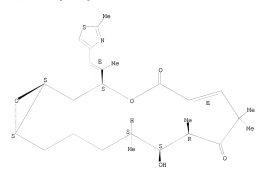
LANGUAGE: English

OTHER SOURCE(S): CASREACT 149:555127

AB A review of the article Dioxirane epoxidn, of alkenes.

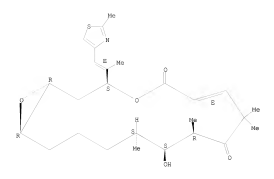
- 193071-71-5P 193071-72-6P RL: SPN (Synthetic preparation); PREP (Preparation) (Dioxirane Epoxidn, of Alkenes)
- RN 193071-71-5 HCAPLUS
- 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, CN 11-hydroxy-8,8,10,12-tetramethyl-3-((1E)-1-methyl-2-(2-methyl-4thiazolvl)ethenvl]-, (1S,3S,6E,10R,11S,12S,16S)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.



- RN 193071-72-6 HCAPLUS
- CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolyl)ethenyl]-, (1R, 3S, 6E, 10R, 11S, 12S, 16R)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.



L7 ANSWER 2 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN ACCESSION NUMBER: 2008:191355 HCAPLUS

DOCUMENT NUMBER: 148:355544

TITLE:

Conformational Preferences of Natural and C3-Modified Epothilones in Aqueous Solution AUTHOR(S):

Erdelyi, Mate; Pfeiffer, Bernhard; Hauenstein, Kurt;

Fohrer, Joerg; Gertsch, Juerg; Altmann, Karl-Heinz; Carlomagno, Teresa NMR-Based Structural Biology, Max-Planck-Institute for CORPORATE SOURCE:

Biophysical Chemistry, Goettingen, D-37077, Germany SOURCE: Journal of Medicinal Chemistry (2008), 51(5),

1469-1473

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER: American Chemical Society DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 148:355544 GT

IT

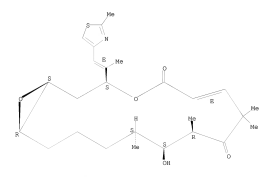
- AR The conformational properties of the microtubule-stabilizing agent epothilone A (I, R = OH, R1 = H) and its 3-deoxy and 3-deoxy-2,3-didehydro derivs. I (R = R1 = H) and I (RR1 = E-bond) have been investigated in aqueous solution by a combination of NMR spectroscopic methods, Monte Carlo conformational searches, and NAMFIS calcns. The tubulin-bound conformation of epothilone A, as previously proposed on the basis of solution NMR data, was found to represent a significant fraction of the ensemble of conformations present for the free ligands in aqueous solution
  - 476623-83-3P RL: PAC (Pharmacological activity); PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent)

(conformational preferences of epothilone A and 3-deoxy derivs. in aqueous solution and antitumor activity)

RN 476623-83-3 HCAPLUS CN

4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolvl)ethenvl]-, (18,38,6E,10R,118,128,16R)- (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.



OS.CITING REF COUNT: 12 THERE ARE 12 CAPLUS RECORDS THAT CITE THIS

RECORD (12 CITINGS)

REFERENCE COUNT: 41 THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 3 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN ACCESSION NUMBER: 2006:396881 HCAPLUS

DOCUMENT NUMBER: 145:240913

TITLE: Prediction of antitumor activity for epothilone

analogues based on 3D molecular descriptors Tan, Ning-Xin; Li, Juan-Qin; Li, Ze-Rong; Li, AUTHOR(S):

Xiang-Yuan

CORPORATE SOURCE: Coll. Chem. Eng., Sichuan Univ., Chengdu, 610065,

Peop. Rep. China

Wuli Huaxue Xuebao (2006), 22(4), 397-402

CODEN: WHXUEU; ISSN: 1000-6818 PUBLISHER: Wuli Huaxue Xuebao Bianjibu Chinese

DOCUMENT TYPE: Journal

LANGUAGE:

In order to predict the antitumor activities of various epothilone analogs, a set of mol. descriptors, including electronic, topol. and geometric descriptors, and mol. shape indexes (K-order moment shape indexes), were calculated to characterize the structural and physicochem. properties for 150 compds. The 30 descriptors selected with genetic algorithm were employed to establish the classification and prediction model of epothilone analogs by using support vector machine (SVM). This SVM system gives a total prediction accuracy of 83.3% by the leave-one-out method and that of 80.6% by the 5-fold cross-validation method. The present study indicates that K-order moment shape indexes are useful for description of configuration isomers, and SVM is a facilitating tool in prediction of antitumor activity of epothilone analogs.

193071-71-5 193071-72-6

RL: DMA (Drug mechanism of action); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(prediction of antitumor activity for epothilone analogs based on 3D mol. descriptors)

RN 193071-71-5 HCAPLUS

CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione,

11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolyl)ethenyl]-, (1S, 3S, 6E, 10R, 11S, 12S, 16S)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

RN 193071-72-6 HCAPLUS

CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4-thiazolyl)ethenyl]-, (1R,3S,6E,10R,11S,12S,16R)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

L7 ANSWER 4 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 2003:757689 HCAPLUS

DOCUMENT NUMBER: 139:276755

TITLE: Preparation of epothilone derivatives for therapeutic

use as anticancer agents

INVENTOR(S): Requeiro-Ren, Alicia; Kim, Soong-Hoon Bristol-Myers Squibb Company, USA PATENT ASSIGNEE(S):

SOURCE: PCT Int. Appl., 47 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

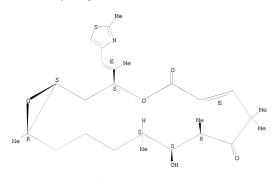
FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

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		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
		PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	TJ,	TM,	TN,	TR,	TT,	TZ,
		UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW						
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,
		KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,
		FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR,
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG
AU	2003	2181	10		A1		2003	0929		AU 2	003-	2181	10		2	0030	311
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PT	1483	251			E		2010	0226		PT 2	003-	7140	96		2	0030	311
	2337				Т3		2010	0421									
RITY	APP	LN.	INFO	. :						US 2							
										WO 2	003-	US75	84	1	W 2	0030	311

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 139:276755 GI

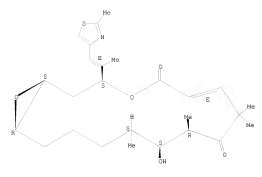
- AB Epothilone derivs., such as I [M = bond, O, NR9, CR10R11; X = O, NH; R1-R4 = H, alkyl; R5 = H, alkyl, cyano; R6 = H, alkyl, aryl, heterocyclyl; R9-R11 = H, OH, alkyl, alkoxy, aryl, cycloalkyl, heterocyclyl], pharmaceutically acceptable salts, solvates or hydrate thereof, were prepared for use as antitumor agents. Thus, epothilone derivative II was prepared
  - from 2,3-dehydro epothilone A, via silvlation of hydroxyl group, potassium cyanide addition, followed by deprotection. The prepared epothilone derivs. were assayed in vitro for their effect on tubulin polymerization and for cytotoxicity against HCT-116 human colon carcinoma cells. Therapeutic compns. containing I or in combination with other therapeutic agents useful in the treatment of cancer or other proliferative diseases are also claimed.
- TТ 226956-21-4 476623-83-3
- RL: RCT (Reactant); RACT (Reactant or reagent)
- (preparation of epothilone derivs, for therapeutic use as anticancer agents) RM 226956-21-4 HCAPLUS
- 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, CN 11-hydroxy-8, 8, 10, 12, 16-pentamethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolv1)ethenv1]-, (1S,3S,6E,10R,11S,12S,16R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.



- RN 476623-83-3 HCAPLUS
- CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolv1)ethenv1]-, (1S,3S,6E,10R,11S,12S,16R)- (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.



OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)

REFERENCE COUNT: THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 5 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN 2002:946278 HCAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER: 138:24591

TITLE:

Preparation of epothilone derivatives for therapeutic use as anti-cancer agents

INVENTOR(S): Requeiro-Ren, Alicia; Borzilleri, Robert M.; Vite,

Gregory D.; Kim, Soong-Hoon PATENT ASSIGNEE(S): Bristol-Myers Squibb Company, USA

SOURCE: PCT Int. Appl., 83 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	ENT I				KIN	D	DATE			APPL	ICAT	ION I	NO.		D	ATE	
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		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,
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		UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZM,	ZW							
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT,	BE,	CH,
		CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG

CA	2449	077			A1		2002	1212	CF	. 2	2002-2	2449	077			20020	514
AU	2002	3098	43		A1		2002	1216	ΑU	1 2	2002-3	3098	43			20020	514
US	2003	0087	888		A1		2003	0508	US	2	2002-	1448	79			20020	514
US	6800	653			B2		2004	1005									
EP	1392	664			A1		2004	0303	EF	2	2002-	7368	67			20020	514
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB, G	R,	IT,	LI,	LU,	NL,	SE	, MC,	PT,
		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY, F	L,	TR						
JP	2004	5328	88		T		2004	1028	JE	2	2003-	5019	91			20020	514
MX	2003	0109	09		A		2004	0217	M	2	2003-	1090	9			20031	127
PRIORITY	APP	LN.	INFO	. :					US	2	2001-2	2954	99P		P	20010	601
									WC	2	2002-0	JS15	397		W	20020	514

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 138:24591

AB Epothilone derivs., such as I [Bl = H, OH, alkoxy, acyloxy, carbamoyl, etc.; W = O, S, NR16; X = O, S, CO, SO, SO2, CH2, CC12, CBr2, NR1, etc.; Rl = H, alkyl; R16 = H, alkyl, aryl, cycloalkyl, heterocyclyl, etc.], were prepared for use as antitumor agents. Thus, aza-epothilone derivative II via a series of synthetic steps which included epoxidn. of epothilone C using 0.0004 M NaZEDTA, F3CCOMe, ZKHSO5.KHSO4.KZSO4 (potassium peroxymonosulfate) and NaHCO3 in MeCN to form epothilone A and 12,13-diepi-epothilone A in 57 and 29% yields, resp., followed by epoxide ring opening/azidation of I2,13-diepi-epothilone A using NaN3 and NH4Cl in EtOH to form the azido-hydroxy derivative in 59% yield, and, finally, formation of II in 62% yield using PPh3 and heating the azido-hydroxy derivative at 60° for 14 h. in THF. The prepared epothilone derivs. were assayed in vitro for their effect on tubulin polymerization and for cytotoxicity

against HCT-116 human colon carcinoma cells.

IT 226956-21-4P

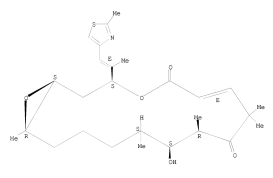
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of epothilone derivs, for therapeutic use as anti-cancer agents)

226956-21-4 HCAPLUS RN

4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hvdroxy-8,8,10,12,16-pentamethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolv1)ethenv1]-, (1S.3S.6E.10R.11S.12S.16R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.



OS.CITING REF COUNT: THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

3

REFERENCE COUNT: THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 6 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 2002:760736 HCAPLUS DOCUMENT NUMBER: 138:95

TITLE:

SAR and pH Stability of Cvano-Substituted Epothilones AUTHOR(S):

Regueiro-Ren, Alicia; Leavitt, Kenneth; Kim, Soong-Hoon; Hoefle, Gerhard; Kiffe, Michael;

Gougoutas, Jack Z.; DiMarco, John D.; Lee, Francis Y. F.; Fairchild, Craig R.; Long, Byron H.; Vite, Gregory

D.

CORPORATE SOURCE: Divisions of Discovery Chemistry Oncology Drug Discovery and Pharmaceutical Development,

Bristol-Myers Squibb Pharmaceutical Research Institute, Princeton, NJ, 08543-4000, USA

SOURCE: Organic Letters (2002), 4(22), 3815-3818 CODEN: ORLEF7; ISSN: 1523-7060

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 138:95

AB 3-Cyano epothilones are the examples of non-hydroxy C-3-substituted analogs. Their tubulin binding affinity and cytotoxicity provide meaningful structure-activity relationship information on the dependence of C-1/C-3 conformation upon activity. 12-Cyano epothilone has improved pH stability over epothilone B, and its activity further supports the hypothesis that C-12 stereochem, is not critical for tubulin affinity.

226956-21-4P 476623-83-3P

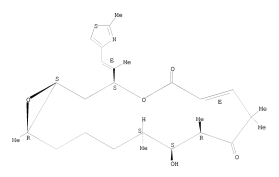
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(pH stability, preparation and structure-activity relationship of cyano-substituted epothilones in human colon carcinoma cells)

RM 226956-21-4 HCAPLUS CN

4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8, 8, 10, 12, 16-pentamethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolv1)ethenv1]-, (15,35,6E,10R,115,125,16R)- (9CI) (CA INDEX NAME)

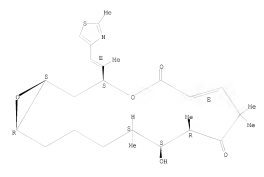
Absolute stereochemistry. Double bond geometry as shown.



RN 476623-83-3 HCAPLUS

CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolv1)ethenv1]-, (1S,3S,6E,10R,11S,12S,16R)- (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.



THERE ARE 26 CAPLUS RECORDS THAT CITE THIS OS.CITING REF COUNT: 26

RECORD (26 CITINGS)

REFERENCE COUNT: 25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 7 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 2002:655116 HCAPLUS

DOCUMENT NUMBER: 137:185358

TITLE: Preparation of epothilone analogs as anticancer agents INVENTOR(S): Nicolaou, Kyriacos C.; He, Yun; Ninkovic, Sacha; Pastor, Joaquin; Roschangar, Frank; Sarabia,

Francisco; Vallberg, Hans; Vourloumis, Dionisios; Winssinger, Nicolas; Yang, Zhen; King, N. Paul;

Finlay, M. Ray PATENT ASSIGNEE(S): The Scripps Research Institute, USA

SOURCE: U.S., 160 pp., Cont.-in-part of U.S. Ser. No.

856,533, abandoned.

CODEN: USXXAM DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

PATENT NO.	KIND DATE	APP	LICATION NO.	DATE
US 6441186	B1 2002	0827 US	1997-923869	19970904
CA 2274833	A1 1998	0618 CA	1997-2274833	19971212
WO 9825929	A1 1998	0618 WO	1997-EP7011	19971212
W: AL, AM, A	I, AU, AZ, BA,	BB, BG, BR	, BY, CA, CH,	CN, CU, CZ, DE,
				KE, KG, KP, KR,
KZ, LC, LI	K, LR, LS, LT,	LU, LV, MD	, MG, MK, MN,	MW, MX, NO, NZ,
PL. PT. RO	D. RU. SD. SE.	SG. SI. SK	. SL. TJ. TM.	TR. TT. UA. UG.

		US,	UZ,	VN,	YU,	ZW											
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SZ,	UG,	ZW,	AT,	BE,	CH,	DE,	DK,	ES,	FI,
		FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,	CI,	CM,
		GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG								
AU	9857	577			A		1998	0703		AU 1	1998~	5757	7		1	9971	212
AU	7465	97			B2		2002	0502									
EP	9446	34			A1		1999	0929	1	EP 1	1997-	9538	80		1	9971	212
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO										
BR	9714	140			A		2000	0229	1	BR I	1997-	1414	0		3	9971	212
	1246						2000	0308		CN 1	1997-	1817	71		1	9971	212
	1134						2004	0114									
JP	2001	5048					2001	0410		JP I	1998-	5262	47		1	9971	212
	6380				B1		2002	0430	1	US 1	1998-	1026	02		1	9980	622
US	2004	0127	432		A1		2004	0701	1	US 2	2003-	7326	98		2	0031	209
US	7173	137			B2		2007	0206									
PRIORIT	Y APP	LN.	INFO	. :					1	US I	1996-	3286	4P		P 1	9961	213
									1	US 1	1997-:	8565	33		B2 1	9970	514
									1	US I	1997-	9238	69		A 1	9970	904
											1997-1				W 1	9971	212
									1	US 1	1999-	3198	85		A3 1	9990	924

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 137:185358 GI

AB Epothilone A, epothilone B, analogs of epothilone and libraries of epothilone analogs of formula I [R1, R2 = H, silyl group, Me, Ac, PhCO, tert-butoxycarbonyl; R3 = H, Me, CHO, (substituted) CO2H, etc.; R4 = heterocyclyl, etc.; X = (CH2)n; n = 1-5) are synthesized. Epothilone A and B are known anticancer agents that derive their anticancer activity by the prevention of mitrois through the induction and stabilization of microtubulin assembly. Several of the analogs are demonstrated to have a superior cytotoxic activities as compared to epothilone A or epothilone B as demonstrated by their enhanced ability to induce the polymerization and stabilization of microtubules. Thus, epothilones A and B are prepared via olefin metathesis and macrocyclization. II was prepared and showed 7% tubulin polymerization

IT 193071-71-5P 193071-72-6P

RL: CPN (Combinatorial preparation); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); CMBI (Combinatorial study); PREP (Preparation); USES (Uses)

(preparation of epothilone analogs as anticancer agents)

RN 193071-71-5 HCAPLUS

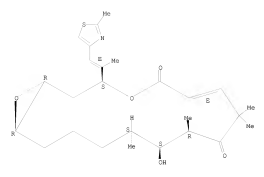
CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8, 8, 10, 12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolvl)ethenvl]-, (18.38,6E,10R,118,128,168)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

RN 193071-72-6 HCAPLUS

CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hvdroxv-8,8,10,12-tetramethvl-3-[(1E)-1-methvl-2-(2-methvl-4thiazolyl)ethenyl]-, (1R,3S,6E,10R,11S,12S,16R)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.



OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD
(2 CITINGS)

REFERENCE COUNT: 31 THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 8 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN ACCESSION NUMBER: 2001:843887 HCAPLUS

DOCUMENT NUMBER: 135:371566

TITLE: Process for reduction of oxiranyl epothilones to

olefinic epothilones

INVENTOR(S): Kim, Soong-hoon; Johnson, James A.
PATENT ASSIGNEE(S): Bristol-Myers Squibb Co., USA

SOURCE: U.S., 10 pp., Cont.-in-part of U.S. Ser. No. 170,581.

DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
US 6320045	B1 20011120	US 1999-316796	19990521
CA 2375029	A1 20001130	CA 2000-2375029	20000515
WO 2000071521	A1 20001130	WO 2000-US13253	20000515
W: AE, AL, AM,	AT, AU, AZ, BA,	BB, BG, BR, BY, CA, CH,	CN, CR, CU,
CZ, DE, DK,	DM, EE, ES, FI,	GB, GD, GE, GH, GM, HR,	HU, ID, IL,
IN, IS, JP,	KE, KG, KP, KR,	KZ, LC, LK, LR, LS, LT,	LU, LV, MA,
MD, MG, MK,	MN, MW, MX, NO,	NZ, PL, PT, RO, RU, SD,	SE, SG, SI,
SK, SL, TJ,	TM, TR, TT, UA,	UG, UZ, VN, YU, ZA, ZW	
RW: GH, GM, KE,	LS, MW, SD, SL,	SZ, TZ, UG, ZW, AT, BE,	CH, CY, DE,
DK, ES, FI,	FR, GB, GR, IE,	IT, LU, MC, NL, PT, SE,	BF, BJ, CF,
CG, CI, CM,	GA, GN, GW, ML,	MR, NE, SN, TD, TG	

EP	1178	968			A1		2002	0213	EF	20	000-9	9307	25			20000	1515
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB, G	R,	IT,	LI,	LU,	NL,	SE	, MC,	PT,
		IE,	SI,	LT,	LV,	FI,	RO										
HU	2002	00146	57		A2		2002	1028	HU	20	002-	1467				20000	515
HU	2002	00146	57		A3		2005	0928									
JP	2003	50039	4		T		2003	0107	JE	20	000-6	6197	78			20000	515
IN	2001	MN011	106		A		2007	0420	IN	1 20	001-1	MN11	06			20010	912
MX	2001	01105	53		A		2002	0722	M	20	001-	1105	3			20011	.030
PRIORIT	APP:	LN. I	NFO	. :					US	19	997-6	6754	9P		P	19971	204
									US	19	998-	8256	3P		P	19980	421
									US	19	998-	1705	81		A2	19981	.013
									US	19	999-	3167	96		A	19990	521
									WC	20	000-	US13	253		W	20000	515

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): CASREACT 135:371566; MARPAT 135:371566 GI

AB This process produced epothilones I (W = O, NR8; Rl-R6 = H, (un) substituted alkyl or aryl and Rl and R2 can be cycloalkyl; R7 = H, (un) substituted alkyl, aryl, cycloalkyl or 4-7 membered heterocyclic N-, O-, or S-containing rings; R8 = H, (un) unsubstituted alkyl, OH, (un) unsubstituted O-alkyl; X = CH2 or XY = CHE-CH; Z = H or OP1 where P1, P2 = H, (un) substituted alkyl, alkanoyl, aroyl, trialkyl (aryl) silyl) from oxiranyl epothilones via the reaction of the oxiranyl moiety with a metal or metal-assisted reagent selected from the group consisting of reactive metallocenes, or (WCl6, n-BuLi). Thus II was prepared in 29% yield in a

multistep reaction from epothilone B via the aminoheptadecenoic acid that cyclized to the oxiranyl azaepothilone intermediate which was reacted with WC16 in THF and n-BuLi in hexane.

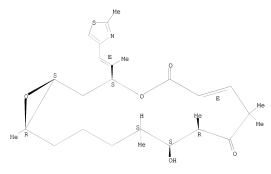
226956-21-4

RL: RCT (Reactant); RACT (Reactant or reagent) (process for reduction of oxiranyl epothilones to olefinic epothilones)

RN 226956-21-4 HCAPLUS

4,17-Dioxabicvclo[14,1,0]heptadec-6-ene-5,9-dione, CN 11-hydroxy-8, 8, 10, 12, 16-pentamethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolyl)ethenyl]-, (15,38,6E,10R,118,128,16R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.



THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD OS.CITING REF COUNT: (2 CITINGS)

REFERENCE COUNT: THERE ARE 88 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 9 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN ACCESSION NUMBER:

2001:137877 HCAPLUS

DOCUMENT NUMBER: 134:335980

TITLE: Comparative molecular field analysis (CoMFA) study of epothilones - tubulin depolymerization inhibitors: pharmacophore development using 3D QSAR methods

AUTHOR(S): Lee, Keun Woo; Briggs, James M.

Department of Biology and Biochemistry, University of Houston, Houston, TX, 77204-5513, USA CORPORATE SOURCE:

Journal of Computer-Aided Molecular Design (2001),

15(1), 41-55

CODEN: JCADEQ; ISSN: 0920-654X

PUBLISHER: Kluwer Academic Publishers

DOCUMENT TYPE: Journal

SOURCE:

LANGUAGE: English

AB A three-dimensional quant, structure-activity relationship (3D OSAR) study has been carried out on epothilones based on comparative mol. field analyses (CoMFA) using a large data set of epothilone analogs, which are potent inhibitors of tubulin depolymn. Microtubules, which are polymers of the  $\alpha/\beta$ -tubulin heterodimer, need to dissociate in order to form the mitotic spindle, a structure required for cell division. A rational pharmacophore searching method using 3D OSAR procedures was carried out and the results for the epothilones are described herein. One-hundred and sixty-six epothilone analogs and their depolymn. inhibition properties with tubulin were used as a training set. Over a thousand mol. field energies were generated and applied to generate the descriptors of QSAR equations. Using a genetic function algorithm (GFA) method, combined with a least square approach, multiple QSAR models were considered during the search for pharmacophore elements. Each GFA run resulted in 100 QSAR models, which were ranked according to their lack of fit (LOF) scores, with a total of 40 GFA runs having been performed. 40 best QSAR equations from each run had adequate fitted correlation coeffs. (R from 0.813 to 0.863) and were of sufficient statistical significance (F value from 7.2 to 10.9). The pharmacophore elements for epothilones were studied by investigating the hit frequency of descriptors (i.e. the sampling probabilities of grid points from the GFA studies) from the set of the 4000 top scoring QSAR equations. By comparing the frequency with which each grid point appeared in the QSAR equations, three candidate regions in the epothilones were proposed to be pharmacophore elements. Two of them are completely compatible with the recent model proposed by Ojima et al. however, one is quite different and is necessary to accurately predict the activities of all 166 epothilone mols. used in our training set. Finally, by visualizing the 35 most probable grid points, it was found that changes related to the C6, C7, C8, C12, S20, and C21 atoms of the epothilones were highly correlated to their activity. 193071-71-5 193071-72-6

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study) (CoMFA study of epothilones - tubulin depolymn. inhibitors:

pharmacophore development using 3D QSAR methods)

RN 193071-71-5 HCAPLUS CN

4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hvdroxv-8, 8, 10, 12-tetramethvl-3-[(1E)-1-methvl-2-(2-methvl-4thiazolyl)ethenyl]-, (1S,3S,6E,10R,11S,12S,16S)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

RN 193071-72-6 HCAPLUS

CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4-thiazolyl)ethenyl]-, (1R,3S,6E,10R,11S,12S,16R)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

OS.CITING REF COUNT: 17 THERE ARE 17 CAPLUS RECORDS THAT CITE THIS

RECORD (17 CITINGS)

REFERENCE COUNT: 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 10 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 2000:842116 HCAPLUS

DOCUMENT NUMBER: 133:362657

TITLE: A process for the reduction of oxiranyl epothilones to

olefinic epothilones INVENTOR(S): Kim, Soong-Hoon; Johnson, James A.

PATENT ASSIGNEE(S): Bristol-Myers Squibb Co., USA

SOURCE: PCT Int. Appl., 19 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

P		ENT 1				KIN	DATE				ICAT				D	ATE	
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							ES,										
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							TT,										
		RW:					SD,										
							GR,							SE,	Br,	BJ,	CF,
11		63200					GW, 2001								- 1	0000	5.2.1
		23750															
		11789					2000										
12	E						ES,										
		14.				LV,		111,	GD,	OI.	11,	шт,	шо,	1411,	JL,	110,	11,
.т	P	20035						0107		TP 2	000-	6197	78		2	0000	515
		20011														0010	
		20010								MX 2	001-	1105	3		2	0011	030
PRIORI	ΤY	APPI	N. :	INFO	. :					US 1	999-	3167	96	- 2	A 1	9990.	521
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										US 1	998-	8256	3P	1	P 1	9980	421
										US 1	998-	1705	81	- 2	A2 1	9981	013
										WO 2	000-1	JS13:	253	1	vi 2	0000	515

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): CASREACT 133:362657; MARPAT 133:362657

GI

12(13)-Olefinic epothilones, such as I and II [R1-6 = H, alkyl, aryl; R1R2 = cycloalkyl; R7 = H, alkyl, aryl, cycloalkyl, heterocyclyl; P1, P2 = H, alkyl, alkanoyl, aroyl, silyl, etc.; W = O, NR8; R8 = H, OH, alkyl], were prepared via reduction of the corresponding 12,13-epoxyepothilones using a metal

or metal-assisted reagent. The metal or metal-assisted reagent was selected from the group consisting of reactive metallocenes, [N2C(CO2Me)2, cat Rh2(OAc)4], [N2C(CO2Me)2, cat[(n-C7H15CO2)2Rh]2], [Zn-Cu, EtOH], [Mg(Hg), MgBr], Cr, [FeCl3, n-BuLi], [TiCl3, LiAlH4], [TiCl4, Zn], [WCl6, LiAlH4], [NDC15, NaAlH4], [VC13, Zn], or [WC16, n-BuLi]. Thus, epothilone A, a 12,13-epoxyepothilone, was reduced using magnesium turnings and titanocene dichloride in THF to give epothilone C, a 12(13)-(Z)-olefin, in 80% vield.

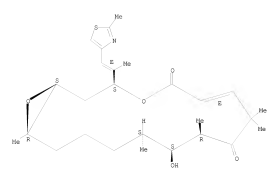
226956-21-4

RL: RCT (Reactant); RACT (Reactant or reagent) (process for the reduction of oxiranyl epothilones to olefinic epothilones)

RN 226956-21-4 HCAPLUS CN

4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12,16-pentamethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolv1)ethenv1]-, (15,3s,6E,10R,11s,12s,16R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.



OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD

(3 CITINGS)

REFERENCE COUNT: THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS 3 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 11 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN ACCESSION NUMBER: 2000:316343 HCAPLUS

Correction of: 1997:528752 DOCUMENT NUMBER: 132:293587

Correction of: 127:149021

TITLE: The Olefin Metathesis Approach to Epothilone A and Its Analogs

AUTHOR(S): Nicolaou, K. C.; He, Y.; Vourloumis, D.; Vallberg, H.; Roschangar, F.; Sarabia, F.; Ninkovic, S.; Yang, Z.;

Trujillo, J. I.

CORPORATE SOURCE: Institute for Chemical Biology, La Jolla, CA, 92037, USA

Journal of the American Chemical Society (1997),

SOURCE: 119(34), 7960-7973 CODEN: JACSAT; ISSN: 0002-7863

PUBLISHER: American Chemical Society

Journal

DOCUMENT TYPE: LANGUAGE:

English GI

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

AB The olefin metathesis approach to epothilone A (I) and several diastereomeric analogs is described. Key building blocks II,

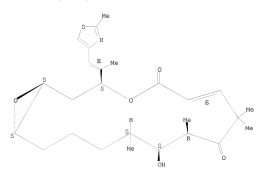
RN

(S)-OHCCH(Me)CH2CH2CH2CH=CH2, and (S)-MeCH2COC(Me)2CH(OSiMe2CMe3)CH2CO2H were constructed in optically active form and were coupled and elaborated to olefin metathesis precursor III (R = SiMe2CMe3) via an aldol reaction and an esterification coupling. Olefin metathesis of compound III (R = SiMe2CMe3), under the catalytic influence of RuCl2(:CHPh)(PCy3)2, furnished cis- and trans-cyclic olefins IV (R = SiMe2CMe3). Epoxidn. of (Z)-IV (R = H) gave I and several analogs, whereas epoxidn. of (E)-IV (R = H) resulted in addnl. epothilones. Similar elaboration of isomeric as well as simpler intermediates resulted in vet another series of epothilone analogs and model systems.

193071-71-5P 193071-72-6P RL: SPN (Synthetic preparation); PREP (Preparation) (synthesis of epothilone A and analogs via olefin metathesis) 193071-71-5 HCAPLUS

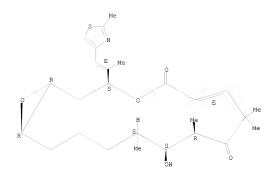
CM 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolyl)ethenyl]-, (1S,3S,6E,10R,11S,12S,16S)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.



193071-72-6 HCAPLUS RN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, CN 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolyl)ethenyl]-, (1R,3S,6E,10R,11S,12S,16R)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.



L7 ANSWER 12 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN ACCESSION NUMBER:

1999:691091 HCAPLUS

DOCUMENT NUMBER: 131:310502

TITLE: synthesis and cytotoxicity of 12,13-modified epothilone derivatives for use in treatment of tumors

or other hyperproliferative cellular disease

INVENTOR(S): Vite, Gregory D.; Kim, Soong-Hoon Kim; Hofle, Gerhard PATENT ASSIGNEE(S): Bristol-Myers Squibb Company, USA

SOURCE: PCT Int. Appl., 89 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA:	TENT :	NO.			KIN	)	DATE			APPL	ICAT	ION I	NO.		D	ATE	
						-									-		
WO	9954	319			A1		1999	1028		WO 1:	999-	US74	75		1	9990	405
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		DK,	EE,	ES,	FI,	GB,	GE,	GH,	GM,	HU,	ID,	IL,	IS,	JP,	KE,	KG,	KP,
		KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,
		NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	UA,
		UG,	UZ,	VN,	YU,	ZA,	ZW										
	RW:	ΑT,	BE,	BF,	ВJ,	CF,	CG,	CH,	CI,	CM,	CY,	DE,	DK,	ES,	FI,	FR,	GA,
			GR,	ΙE,	ΙT,	LU,	MC,	ML,	MR,	ΝE,	NL,	PT,	SE,	SN,	TD,	TG	
US	6380	395			В1		2002	0430		US 1	999-	2801	92		1	9990	329
US	6399	638			В1		2002	0604		US 1	999-	2801	91		1	9990	329
CA	2329	181			A1		1999	1028		CA 1:	999-	2329	181		1:	9990	405
AU	9934	716			A		1999	1108		AU 1	999-	3471	6		13	9990	405
AU	7485	26			B2		2002	0606									
BR	9909	795			A		2000	1226		BR 1	999-	9795			1	9990	405

TR	2000003	036		T2	2001	0122	TR	2000-	3036				19990	405
EP	1073648			A1	2001	0207	EP	1999-	9163	83			19990	405
EP	1073648			B1	2006									
	R: AT		CII		DK, ES,		CP C	р тт	TT	TIT	MI	CE	MC	DT
		, FI,		DE,	DIC, ES,	LIC	GD, G	, 11,	шт,	шо,	ML,	JE	, PEC,	LI,
			CI						_					
JP	2002512	239		T	2002	0423	JP	2000-	5446	58			19990	405
CN	1142923			С	2004	0324	CN	1999-	8052	66			19990	405
EP	1589017			A2	2005	1026	EP	2005-	1523	6			19990	405
EP	1589017			A3	2009	0422								
	R: AT		CH.		DK, ES,		GB. GI	R. TT.	T.T.	T.U.	NI	SE	. MC.	PT.
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AT	340177			T	2006	1015	AT	1999-	9163	83			19990	405
PT	1073648	:		E	2006	1229	PT	1999-	9163	83			19990	405
ES	2273484			Т3	2007	0501	ES	1999-	9163	83			19990	405
PT	1073647			E	2009	0717	PT	1999-	9152	73			19990	405
	2327803			T3		1103		1999-					19990	
	2000010			A		0419		2000-					20001	
				**	2001	0113								
PRIORITY	APPLN.	TMF.C	. :					1998-					19980	
							EP	1999-	9163	83			19990	
							WO	1999-	US74	75		W	19990	405

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 131:310502

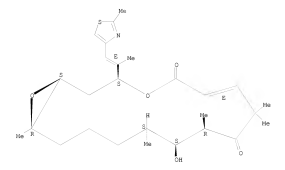
GI

AB Synthesis and cytotoxicity of 12,13-modified epothilone derivs.(I) [R1 = H, (un)substituted alkyl; R2 = H if bond double or βOH if bond single; Y = O, NH; X = O, (un)substituted NH, OCH2, 2-methylthiazolo, S, (un)substituted CH2] is presented. Thus, I (R1 = H, X = NH, R2 = βOH, Y = O) (II) is prepared by epoxidn. of epothilone C followed by azidation and reductive imination. I are useful in treatment of tumors or other hyperproliferative cellular disease and show IC50 of 0.01-1000 nM in cell proliferation tests.

Ι

- RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
  - (synthesis and cytotoxicity of 12,13-modified epothilone derivs. for use in treatment of tumors or other hyperproliferative cellular disease)
- RN 226956-21-4 HCAPLUS
- CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 1-hydroxy-8,8,10,12,16-pentamethyl-3-[(1E)-1-methyl-2-(2-methyl-4-thiazolyl)ethenyl]-, (1S,3S,6E,10R,11S,12S,16R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.



OS.CITING REF COUNT: (5 CITINGS)

THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 13 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1999:375551 HCAPLUS

DOCUMENT NUMBER: 131:31830

TITLE: A process for the reduction of oxiranyl epothilones to

olefinic epothilones Kim, Soong-Hoon; Johnson, James A.

PATENT ASSIGNEE(S): Bristol-Myers Squibb Company, USA

SOURCE: PCT Int. Appl., 19 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

INVENTOR(S):

P	ATENT	NO.			KIN	D	DATE			APPL.	ICAT	ION :	NO.		D	ATE		
-						_									-			
W	0 9928	324			A1		1999	0610		WO 1	998-	US25	464		1	9981	201	
	W:	AL,	AM,	ΑT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,	
		DK,	EE,	ES,	FI,	GB,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IS,	JP,	KE,	KG,	
		KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	
		NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	
		UA,	UG,	UZ,	VN,	YU,	zw											
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SZ,	UG,	ZW,	ΑT,	BE,	CH,	CY,	DE,	DK,	ES,	
		FT	FR	GB	GR	TE	TT	T.II	MC	MT.	PT	SE	BF	P.T	CE	CG	CT	

			CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TE	, TG							
	CA	2311	929			A1	A1 19990610				CA 1998-2311929						19981201		
	ΑU	9915	408			A 19990616				AU 1999-15408						19981201			
	ΑU	7385	76			B2 20010920													
1	EP	1042	327			A1 20001011				EP 1998-959652						19981201			
1	EΡ	1042	327		B1 20030917														
				BE.	CH.					GB.	GF	IT.	LI.	LU.	NL.	SE.	MC.	PT.	
			IE.	FI															
1	HU 2001000582						A2 20010928 HU 2001-582									19981201			
1	HU 2001000582							2003	0328										
	JP 2001525324							2001	1211	J	P	2000-	5232	16		1	9981	201	
	JΡ	4434	484			B2		2010	0317										
	IL	1355	90			A		2003	0917	I	L	1998-	1355	90		1	9981	201	
AT 250066						т		2003	1015	A	т	1998-	9596	52		1	9981	201	
1	ES	2207	015			Т3		2004	0516	Ε	S	1998-	9596	52		1	9981	201	
	TW	2214	69			В		2004	1001	Т	W	1998-	8711	9880		1	9981	201	
1		A1		2004	0514	H	K	2000-	1078	69		2	0001	207					
PRIOR	IΤ	APP	LN.	INFO	. :					U	S	1997-	6754	9P		P 1	9971	204	
										U	S	1998-	8256	3P		P 1	9980	421	
										W	0	1998-	US25	464		W 1	9981	201	
OTHER SOURCE(S): GI							REAC	T 13	1:31	830;	MA	RPAT	131:	3183	0				

AB The olefinic epothilones I and II (X = 0, NR8; Z = bond, Rl-R6 = H, alkyl, substituted alkyl, aryl, RlR2 may be a cycloalkyl; R7 = H, alkyl, substituted alkyl, aryl, cycloalkyl, heterocyclo; R8 = H, alkyl, substituted alkyl, OH, alkoxy, substituted alkoxy; Pl, P2 = H, alkyl, substituted alkyl, alkanoyl, substituted alkoxy; Pl, P2 = H, alkyl, substituted alkyl, alkanoyl, substituted aroyl, trialkylsilyl, aryldialkylsilyl, diarylalkylsilyl, triarylsilyl) were prepared by reduction of the oxiranyl epothilones I and II (Z = 0) with a

ΙI

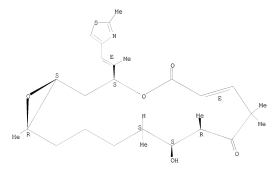
metal or metal assisted reagents, e.g. metallocenes, WC14-BuLi, VC13-Zn, TiCl3-LiAlH4. Thus, epothilone A was treated with Mg and

bis(cyclopentadienyl)titanium dichloride in THF to give 80% epothilone C.
II 226956-21-4
RL: RCT (Reactant); RACT (Reactant or reagent)

(process for reduction of oxiranyl epothilones to olefinic epothilones)
226956-21-4 HCAPLUS

RN 226956-21-4 HCAPLUS
0 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione,
11-hydroxy-8,8,10,12,16-pentamethyl-3-[(15)-1-methyl-2-(2-methyl-4-thiazolyl)ethenyl]-, (18,38,6E,10R,118,125,16B)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.



OS.CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD
(4 CITINGS)

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 14 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1998:760825 HCAPLUS

DOCUMENT NUMBER: 130:95406

TITLE: Oxidative and reductive transformations of epothilone

AUTHOR(S): Sefkow, Michael; Kiffe, Michael; Schummer, Dietmar; Hofle, Gerhard

CORPORATE SOURCE: Gesellschaft fur Biotechnologische Forschung mbH, Abt, Naturstoffchemie, Braunschweig, D-38124, Germany

SOURCE: Bioorganic & Medicinal Chemistry Letters (1998),

8(21), 3025-3030 CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal

10591921z.trn 06/04/2010

Page 36

LANGUAGE: English

OTHER SOURCE(S): CASREACT 130:95406

AB The C7 hydroxy group of cytotoxic epothilone A was selectively oxidized using PDC. A selective oxidation of the C3 hydroxy group was accomplished with Me25/(PhCO2)2 after in situ protection of C7-OH. Reduction of epothilone A or of a C5, C7 dioxo derivative with NaBH4 proceeded at the C5 carbonyl group. Oxidation and hydrogenation of the C16-C17 double bond proved to be difficult but it was easily cleaved with ozone and the resulting keto derivative was transformed to epothilone analoso with different side chains.

IT 219557-03-6P

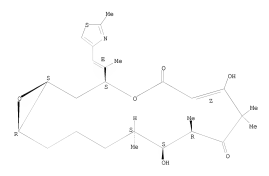
RI: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(oxidative and reductive transformations of epothilone A)

RN 219557-03-6 HCAPLUS CN 4.17-Dioxabicvclo[14

4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione,
7,11-dihydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolyl)ethenyl]-, (18,38,67,10R,118,128,16R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.



OS.CITING REF COUNT: 26 THERE ARE 26 CAPLUS RECORDS THAT CITE THIS

RECORD (26 CITINGS)

REFERENCE COUNT: 40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 15 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1998:405952 HCAPLUS

DOCUMENT NUMBER: 129:81625

ORIGINAL REFERENCE NO.: 129:16853a,16856a

TITLE: Preparation of epothilone analogs as anticancer agents
INVENTOR(S): Nicolaou, Costa Kyriacos; He, Yun; Ninkovic, Sacha;

Page 37

Pastor, Joaquin; Roschangar, Frank; Sarabia, Francisco; Vallberg, Hans; Vourloumis, Dionisios; Winssinger, Nicolas; Yang, Zhen; King, Nigel Paul; et

PATENT ASSIGNEE(S): SOURCE:

Novartis A.-G., Switz.; Scripps Research Institute PCT Int. Appl., 213 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

	PATENT NO.																		
												1997-							
		W:										, BY,							
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			PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK	, SL,	TJ,	TM,	TR,	TT,	UA,	UG,	
			US,	UZ,	VN,	YU,	ZW												
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								SN,											
Ţ	JS	6441	186			B1		2002	0827		US	1997-	9238	69		1	19970	904	
(	CA 2274833					A1 19980618				US 1997-923869 CA 1997-2274833					19971212				
2	AU 9857577								AU 1998-57577					19971212					
2	ΑU	74659	97			B2		2002	0502										
1	ΞP	94463	34			A1		1999	0929		EP	1997-	9538	08		1	19971	212	
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	PT,	
			IE,	SI,	LT,	LV,	FI,	RO											
3	ЗR	97143	140			A		2000	0229		BR	1997- 1998- 1999-	1414	0		1	19971	212	
	JΡ	20015	5048	56		T		2001	0410		JP	1998-	5262	47		1	19971	212	
Ţ	JS	6660	758			B1		2003	1209		US	1999-	3198	85		1	19990	924	
Ţ	JS	20040	127	432		A1		2004	0701		US	2003-	7326	98		2	20031	209	
Ţ	JS	71733	137			B2		2007	0206			2003-							
PRIOR:											US	1996-	3286	4P		P 1	19961	213	
												1997-							
											US	1997-	9238	69		A2 1	19970	904	
											WO	1997-	EP70	11		W 1	19971	212	
											US	1999-	3198	85		A3 1	19990	924	
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 129:81625

GI

- AB Epothilone A, epothilone B, analogs of epothilone and libraries of epothilone analogs of formula I [X = (CH2)n, n = 1-5; R 1 = OH, OMe, absent; R2, R3 = H, CH2, Me; R4 = H, Me, protecting group; R5 = H, Me, CHO, (substituted) CO2H, etc.; R6 = O, CH2, absent; R7 = thiazolealkyl, etc.] are synthesized. Epothilone A and B are known anticancer agents that derive their anticancer activity by the prevention of mitosis through the induction and stabilization of microtubulin assembly. Several of the analogs are demonstrated to have a superior cytotoxic activity as compared to epothilone A or epothilone B as demonstrated by their enhanced ability to induce the polymerization and stabilization of microtubules. Thus, II was prepared and was shown to induce tubulin polymerization at 94% relative to GTP, and
- inhibit carcinoma cell growth.
- IT 193071-71-5P 193071-72-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of epothilone analogs as anticancer agents)

RN 193071-71-5 HCAPLUS

- CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione,
  - 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4-thiazolyl)ethenyl]-, (1S,3S,6E,10R,11S,12S,16S)- (CA INDEX NAME)

RN 193071-72-6 HCAPLUS

CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4-thiazolyl)ethenyl]-, (1R,38,6E,10R,11S,12S,16R)- (CA INDEX NAME)

SOURCE:

OS.CITING REF COUNT: 28 THERE ARE 28 CAPLUS RECORDS THAT CITE THIS

RECORD (28 CITINGS)

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 4

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 16 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1997:714315 HCAPLUS

DOCUMENT NUMBER: 128:3560

ORIGINAL REFERENCE NO.: 128:771a

TITLE: Designed epothilones: combinatorial synthesis, tubulin assembly properties, and cytotoxic action against

taxol-resistant tumor cells

AUTHOR(S): Nicolaou, K. C.; Vourloumis, Dionisios; Li, Tianhu;

Pastor, Joaquin; Winssinger, Nicolas; He, Yun;

Ninkovic, Sacha; Sarabia, Francisco; Vallberg, Hans;

Roschangar, Frank; King, N. Paul; Finlay, M. Ray V.;

Giannakakou, Pareskevi; Verdier-Pinard, Pascal; Hamel, Ernest

CORPORATE SOURCE:

Department of Chemistry and The Skaggs Institute for Chemical Biology, The Scripps Research Institute, La

Jolla, CA, 92037, USA

Angewandte Chemie, International Edition in English

(1997), 36(19), 2097-2103

CODEN: ACIEAY; ISSN: 0570-0833

PUBLISHER: Wiley-VCH DOCUMENT TYPE: Journal

LANGUAGE: English

The title work demonstrates the power of interfacing combinatorial chemical with chemical biol. as facilitated by solid-phase synthesis, radiofrequency encoded combinatorial chemical and modern biol. assays. A library of 112 epothilones were prepared by solid-phase synthesis, their structure activity relationships measured by tubulin binding assay and some tested for

inhibition of carcinoma cell growth.

193071-71-5P 193071-72-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(combinatorial synthesis of epothilone library, tubulin assembly properties, and cytotoxic action against taxol-resistant tumor cells)

RN 193071-71-5 HCAPLUS

CM 4,17-Dioxabicvclo[14.1.0]heptadec-6-ene-5,9-dione,

11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolyl)ethenyl]-, (1S,3S,6E,10R,11S,12S,16S)- (CA INDEX NAME)

RN 193071-72-6 HCAPLUS

CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4-thiazolyl)ethenyl]-, (1R,3S,6E,10R,11S,12S,16R)- (CA INDEX NAME)

OS.CITING REF COUNT: 200 THERE ARE 200 CAPLUS RECORDS THAT CITE THIS

RECORD (204 CITINGS)

REFERENCE COUNT: 62 THERE ARE 62 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 17 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1997:528752 HCAPLUS DOCUMENT NUMBER: 127:149021

ORIGINAL REFERENCE NO.: 127:28789a,28792a

TITLE: The Olefin Metathesis Approach to Epothilone A and Its

Analogs AUTHOR(S):

Nicolaou, K. C.; He, Y.; Vourloumis, D.; Vallberg, H.; Roschangar, F.; Sarabia, F.; S.Ninkovic; Yang, Z.;

Trujillo, J. I.

CORPORATE SOURCE: Department of Chemistry and The Skaggs, Institute for Chemical Biology, La Jolla, CA, 92037, USA

SOURCE: Journal of the American Chemical Society (1997),

119(34), 7960-7973

CODEN: JACSAT; ISSN: 0002-7863 PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 127:149021

For diagram(s), see printed CA Issue.

The olefin metathesis approach to epothilone A (I) and several diastereomeric analogs is described. Key building blocks II,

(S)-OHCCH(Me)CH2CH2CH2CH2CH2CH2, and (S)-MeCH2COC(Me)2CH(OSiMe2CMe3)CH2CO2H were constructed in optically active form and were coupled and elaborated to olefin metathesis precursor III (R = SiMe2CMe3) via an aldol reaction and an esterification coupling. Olefin metathesis of compound III (R = SiMe2CMe3), under the catalytic influence of RuCl2(:CHPh)(PCy3)2, furnished cis- and trans-cyclic olefins IV (R = SiMe2CMe3). Epoxidn. of

(Z)-IV (R = H) gave I and several analogs, whereas epoxidn. of (E)-IV (R = H) resulted in addnl. epothilones. Similar elaboration of isomeric as well as simpler intermediates resulted in yet another series of epothilone analogs and model systems.

193071-71-5P 193071-72-6P

RL: SPN (Synthetic preparation); PREP (Preparation)

(synthesis of epothilone A and analogs via olefin metathesis)

RN 193071-71-5 HCAPLUS CM

4.17-Dioxabicvclo[14.1.0]heptadec-6-ene-5.9-dione. 11-hydroxy-8,8,10,12-tetramethyl-3-[(1E)-1-methyl-2-(2-methyl-4thiazolyl)ethenyl]-, (1S,3S,6E,10R,11S,12S,16S)- (CA INDEX NAME)

RN 193071-72-6 HCAPLUS

13001-12 of maximum (4,17-Dioxabicyclo|14.1.0]heptadec-6-ene-5,9-dione,
11-hydroxy-8,8,10,12-teramethyl-3-|(1E)-1-methyl-2-(2-methyl-4-thiazolyl)ethenyl]-,(1R,3S,6E,10R,11S,12S,16E)- (CA INDEX NAME) CN

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

L7 ANSWER 18 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1997:456769 HCAPLUS DOCUMENT NUMBER: 127:50474

ORIGINAL REFERENCE NO.: 127:9629a

Preparation of epothilone derivatives as agrochemicals TITLE: and pharmaceuticals

INVENTOR(S): Hoefle, Gerhard; Kiffe, Michael

PATENT ASSIGNEE(S): Gesellschaft fuer Biotechnologische Forschung Mbh

(Gbf), Germany SOURCE: Ger. Offen., 9 pp.

CODEN: GWXXBX DOCUMENT TYPE: Patent

German LANGUAGE: FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PA:	TENT NO.			KIN	DATE			APE	PLI	CAT	ION	NO.		D	ATE		
DE WO	19542986 9719086 W: JP,	IIS		A1 A1	1997 1997	0522 0529		DE WO	19 19	95- 96-l	1954 EP50	2986 80		1:	9951 9961	117 118	
	RW: AT,	BE,	CH,	DE,	DK, ES,	FI,	FR,	GE	3,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE
ΕP	873341 873341			B1	2003	0910											
	R: AT, IE,	mr.															
EP EP	903348 903348 903348			A1 B1	1999 2002	0324 0605		EP	19	98-	1215	23		1	9961	118	
ΕP	903348			B2	2008	0827											
	R: AT.	BE.	CH.	DE.	DK. ES.	FR.	GB.	GF	₹.	IT.	LI.	LU.	NL.	SE.	MC.	PT.	
	IE, 20005007 4183099 1186606	FI															
JP	20005007	57		т	2000	0125		JP	19	97-	5193	81		1	9961	118	
JP	4183099			B2	2008	1119											
EP	1186606			A1	2002	0313		EΡ	20	01-	1273	52		1	9961	118	
	R: AT, IE, 218556 903348 2178093 249463 873341 261961 2206607 1440973 1440973 R: AT	BE,	CH,	DE,	DK, ES,	FR,	GB,	GF	₹,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
ΑT	218556			T	2002	0615		AΤ	19	98-	1215	23		1	9961	118	
PΤ	903348			E	2002	1129		PΤ	19	98-	1215	23		1:	9961	118	
ES	2178093			Т3	2002	1216		ES	19	98-	1215	23		1	9961	118	
ΑT	249463			T	2003	0915		AΤ	19	96-	9390	97		1	9961	118	
PΤ	873341			E	2004	0227		PΤ	19	96-	9390	97		1	9961	118	
ΑT	261961			T	2004	0415		AΤ	20	01-	1273	52		15	9961	118	
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ΕP	1440973			A2	2004	0728		EΡ	20	04-	5011			13	9961	118	
ΕP	1440973			A3	2004	1020											
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PT	1186606			E	2004	0831		PΤ	20	01-	1273	52		1	9961	118	
ES	2218328			Т3	2004	1116		ES	20	01-	1273	52		15	9961	118	
US	6288237			В1	2001	0911		US	19	98-	7705	5		1	9980	803	
US	1186606 2218328 6288237 20010034 6613912 20040087	452		A1 B2	2001	1025		US	20	01-	8361	34		2	0010	416	
US	20040087	634		A1	2004	0506		US	20	03-	6027	70		2	0030	625	

US 6831076 B2 20041214

PRIORITY APPLN. INFO.: DE 1995-19542986 A 19951117
DE 1996-19639456 A 19960925

EP 1996-939097 A3 19961118 EP 2001-127352 A3 19961118 WO 1996-EP5080 W 19961118 US 1998-77055 A3 19980803 US 2001-836134 A3 20010416

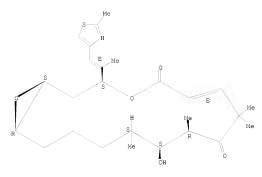
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 127:50474

GI

- AB The title compds., e.g., I [R = H, Cl-4 alkyl, Rl, R2 = H, Cl-6 alkyl, Cl-6 acyl, benzoyl, Cl-4 trialkylsilyl, benzyl, Ph, Cl-6 alkoxy, C6 alkyl-, hydroxy-, and halo-substituted benzyl or phenyl; X, Y = halo, OH, acyloxy, alkoxy, benzoyloxy], useful as agrochems. and pharmaceuticals (no data), are prepared Thus, epothilone A in acetone containing trifluoroacetic acid was heated overnight at 50° and the reaction mixture was adjusted to pH 7 with 1 M phosphate buffer to give 2 isomers, each in 19% yield.
- IT 191105-88-1P RL: AGR (Agricultural use); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of epothilone derivs. as acrochems. and pharmaceuticals)
- RN 191105-88-1 HCAPLUS
- CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione,
  11-hydroxy-8,8,10,12-tetramethyl-3-[1-methyl-2-(2-methyl-4thiazolyl)ethenyl]-, [1R\*,3R\*(E),6E,10S\*,11R\*,12R\*,16S\*]- (9CI) (CA INDEX NAME)

Relative stereochemistry. Double bond geometry as shown.



OS.CITING REF COUNT: THERE ARE 9 CAPLUS RECORDS THAT CITE THIS RECORD (12 CITINGS)

L7 ANSWER 19 OF 19 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 1997:443365 HCAPLUS

DOCUMENT NUMBER: 127:81289

ORIGINAL REFERENCE NO.: 127:15585a,15588a

TITLE: Preparation of epothilone derivatives as agrochemicals

and pharmaceuticals

INVENTOR(S): Hofle, Gerhard; Kiffe, Michael

PATENT ASSIGNEE(S): Gesellschaft Fur Biotechnologische Forschung Mbh (Gbf), Germany; Hofle, Gerhard; Kiffe, Michael

PCT Int. Appl., 38 pp.

SOURCE: CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PA:	TENT	NO.			KIN	D	DATE		API	PLICAT	CION	NO.		D.	ATE		
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WO	9719	086			A1		1997	0529	WO	1996-	-EP50	80		1	9961	118	
	₩:	JP,	US														
	RW:	AT,	BE,	CH,	DE,	DK,	ES,	FI,	FR, G	3, GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE
DE	1954	2986			A1		1997	0522	DE	1995-	-1954	2986		1	9951	117	
DE	1963	9456			A1		1998	0326	DE	1996-	-1963	9456		1	9960	925	
EP	8733	41			A1		1998	1028	EP	1996-	-9390	97		1	9961	118	
EP	8733	41			B1		2003	0910									
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB, GI	R, IT,	LI,	LU,	NL,	SE,	MC,	PT,	
		IE,	FI														
JP	2000	5007	57		T		2000	0125	JP	1997-	-5193	81		1	9961	118	
JP	4183	099			B2		2008	1119									

10591921z.trn 06/04/2010 Page 47

AT 249463 US 6288237	T B1	20030915		1996-939097 1998-77055		19961118 19980803
US 20040087634 US 6831076	A1 B2	20040506		2003-602770		20030625
PRIORITY APPLN. INFO.:			DE WO US	1995-19542986 1996-19639456 1996-EP5080 1998-77055 2001-836134	A W A3	19951117 19960925 19961118 19980803 20010416

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 127:81289

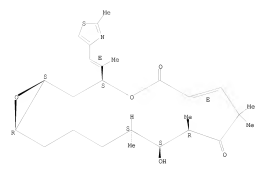
AB The title compds, e.g., I [R = H, C1-4 alkyl, Rl, R2 = H, C1-6 alkyl, C1-6 acyl, benzoyl, C1-4 trialkylsilyl, benzyl, Ph, C1-6 alkoxy, C6 alkyl-, hydroxy-, and halo-substituted benzyl or phenyl, X, Y = H, halo, pseudohalo, OH, acyloxy, alkoxy, benzoyloxy; or YZ = O, bond, however, I may not be epothilone A or Bl, useful as agrochems. and pharmaceuticals (no data), are prepared Thus, epothilone A in acetone containing trifluoroacetic acid was heated overnight at 50° and the reaction mixture was adjusted to pH 7 with 1 M phosphate buffer to give 2 isomers, each in 19% yield.

IT 191105-88-1P

Ι

- RL: AGR (Agricultural use); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
- (preparation of epothilone derivs. as agrochems. and pharmaceuticals)
  RN 191105-88-1 HCAPLUS
- CN 4,17-Dioxabicyclo[14.1.0]heptadec-6-ene-5,9-dione, 11-hydroxy-8,8,10,12-tetramethyl-3-[1-methyl-2-(2-methyl-4thiazolyl)ethenyl]-, [1R\*,3R\*(E),6E,10S\*,11R\*,12R\*,16S\*]- (9CI) (CA INDEX NAME)

Relative stereochemistry. Double bond geometry as shown.



THERE ARE 27 CAPLUS RECORDS THAT CITE THIS OS.CITING REF COUNT: 27 RECORD (31 CITINGS)

REFERENCE COUNT: THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS 1 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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